

Docket No.: VANM213.001AUS

October 12, 2005

Page 1 of 1

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Please Direct All Correspondence to Customer Number **20995**

PETITION TRANSMITTAL LETTER

Applicant : Remacle et al.
App. No : 09/817,014
Filed : March 23, 2001
For : IDENTIFICATION OF BIOLOGICAL
(MICRO)ORGANISMS BY DETECTION
OF THEIR HOMOLOGOUS
NUCLEOTIDE SEQUENCES ON
ARRAYS
Examiner : Calamita, Heather
Art Unit : 1637

CERTIFICATE OF MAILING

I hereby certify that this correspondence and all marked attachments are being deposited with the United States Postal Service as first-class mail in an envelope addressed to: Mail Stop: Petitions, Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450, on

October 12, 2005

(Date)

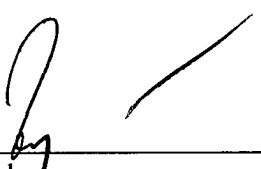
Marina L. Gordey, Reg. No. 52,950

Mail Stop Petitions
Commissioner for Patents
P.O. Box 1450
Alexandria, VA 22313-1450

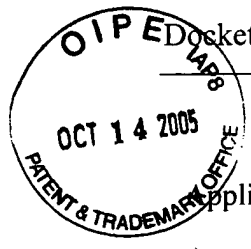
Dear Sir:

Enclosed for filing in the above-identified application are:

- (X) Petition under 37 CFR 1.181(a) to Withdraw Holding of Abandonment Based on Evidence that a Reply was Timely Mailed or Filed.
- (X) Copy of PTO stamped postcard and response filed September 19, 2005.
- (X) The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment, to Account No. 11-1410.
- (X) Return prepaid postcard.



Marina L. Gordey
Registration No. 52,950
Agent of Record
Customer No. 20,995
(805) 547-5580



Docket No.: VANM213.001AUS

Customer No. 20,995

PETITION

Applicant : Remacle et al.
App. No : 09/817,064
Filed : March 23, 2001
For : IDENTIFICATION OF BIOLOGICAL
(MICRO)ORGANISMS BY DETECTION OF THEIR
HOMOLOGOUS NUCLEOTIDE SEQUENCES ON
ARRAYS
Examiner : Calamita, Heather
Art Unit : 1637

**PETITION UNDER 37 CFR 1.181(a) TO WITHDRAW HOLDING OF ABANDONMENT
BASED ON EVIDENCE THAT A REPLY WAS TIMELY MAILED OR FILED**

Mail Stop: Petitions
Commissioner for Patents
P.O. Box 1450
Alexandria, VA 22313-1450

Dear Sir:

We are in receipt of a Notice of Abandonment for the above-identified application, which was mailed on September 30, 2005.

We request withdrawing holding of Abandonment since we timely filed a response to the March 18, 2005 Office Action before the three-month extension of time date on Monday, September 19, 2005.

Enclosed is a copy of the PTO stamped postcard and response filed on September 19, 2005 and date-stamped on September 21, 2005.

Respectfully submitted,

KNOBBE, MARTENS, OLSON & BEAR, LLP

Dated: October 12, 2005

By: _____

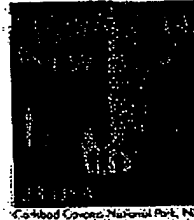
Marina L. Gordey
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Krabbe, Martens, Olson & Bear LLP



LAW OFFICES OF
KNOBBE, MARTENS, OLSON & BEAR, LLP

INTELLECTUAL PROPERTY LAW

2040 Main Street
14th Floor
Irvine, CA 92614

**UTILITY/DESIGN PATENT
(RCE)**

Date: 7-19-05

Date of Office Action: 3-18-05

Notice of Appeal Filed:

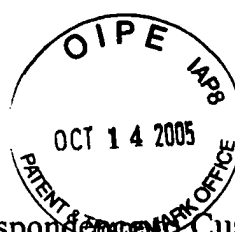
Rec'd in the USPTO on the date stamped hereon via Certificate of Mail:
Docket #: VA1UM313.001AVUS Applicant: REMACLE ET AL.
Title: DENTAL IMPLANT OF BIOLOGICAL (MICRO)ORGANISMS...
App No.: 09/181,704 Filed: 3-23-01
VERIFIED BY: Asst: UKOOPM Atty: MXG QC:

- Request for Continued Examination
- Amendment in 15 pgs
- 3 Month Extension Requested
- Request for Suspension for 3 Months
- Information Disclosure Statement with PTO-1449 w/ 3 Ref(s)
- Terminal Disclaimer in _____ pgs
- Sequence Submission Statement
- Sequence Listing in _____ pgs
- _____ copies of CRF Containing Seq. List
- Return Prepaid Postcard

- Req for Drawing Changes/Corrections
- _____ sheets of RED-Lined Drawings
- Affidavit(s)/Declaration(s)
- \$ 1,940 Check for Filing Fees
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SCP-B-1-2005

(7/14/03)



September 19, 2005
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H.R.C. 2205

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REQUEST FOR CONTINUED EXAMINATION

Applicant	:	Remacle et al.
App. No	:	09/817,014
Filed	:	March 23, 2001
For	:	IDENTIFICATION OF BIOLOGICAL (MICRO)ORGANISMS BY DETECTION OF THEIR HOMOLOGOUS NUCLEOTIDE SEQUENCES ON ARRAYS
Examiner	:	Calamita, Heather
Art Unit	:	1637

CERTIFICATE OF MAILING

I hereby certify that this correspondence and all marked attachments are being deposited with the United States Postal Service as first-class mail in an envelope addressed to: Mail Stop: RCE, Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450, on

September 19, 2005

(Date)

[Signature]
Marina L. Gordey, Reg. No. 52,950

Mail Stop RCE
Commissioner for Patents
P.O. Box 1450
Alexandria, VA 22313-1450

Dear Sir:

This Request for Continued Examination (RCE) is being made as follows:

1. Submission Required under 37 CFR 1.114:

- Enclosed:
- Amendment/Reply in 15 pages.
- Information Disclosure Statement and PTO/SB/08 Equivalent in 2 total pages (IDS and PTO/SB/08).
- (3) references enclosed.
- Return Postcard.

2. Miscellaneous:

- Suspension of action on the above-identified application is requested under 37 CFR § 1.103(c) for a period of 3 months. (Period of suspension shall not exceed three months).

09/22/2005 MBIZUNES 00000024 111410 09817014

01 FC:1801 790.00 OP

09/22/2005 MBIZUNES 00000024 111410 09817014

02 FC:1253	1020.00 OP
03 FC:1463	70.00 DA

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3. Fees:

FEE CALCULATION				
FEE TYPE		FEE CODE	CALCULATION	TOTAL
RCE Fee		1801 (\$790)		\$790
Suspension of Action		1463 (\$130)		\$130
Total Claims	36 - 39 = 0	1202 (\$50)	0 x 50 =	\$0
Independent Claims	2 - 3 = 0	1201 (\$200)	0 x 200 =	\$0
Multiple Claim		1203 (\$360)		\$0
3 Month Extension		1253 (\$1,020)		\$1,020
			TOTAL FEE DUE	\$1,940

(X) An extension of time is hereby requested by payment of the appropriate fee indicated above.

4. Payment:

(X) Check in the amount of \$1,940 to cover the above fees.

Please charge any additional fees, including any fees for additional extension of time, or credit overpayment to Deposit Account No. 11-1410.

Respectfully submitted,
KNOBBE MARTENS OLSON & BEAR LLP

Marina E. Gordey
Registration No. 52,950
Agent of Record
Customer No. 20,995
(805) 547-5580

1939366:vr
091905



VANM213.001AUS

PATENT

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant	:	Remacle et al.
Appl. No.	:	09/817,014
Filed	:	March 23, 2001
For	:	IDENTIFICATION OF BIOLOGICAL (MICRO)ORGANISMS BY DETECTION OF THEIR HOMOLOGOUS NUCLEOTIDE SEQUENCES ON ARRAYS
Examiner	:	Calamita, Heather
Group Art Unit	:	1637

AMENDMENT AND RESPONSE TO OFFICE ACTION WITH RCE

Mail Stop Amendment
Commissioner for Patents
P.O. Box 1450
Alexandria, VA 22313-1450

Dear Sir:

In response to the Office Action mailed March 18, 2005 in the above-identified application, Applicant responds as follows:

Amendments to the Claims are reflected in the listing of claims which begins on page 2 of this paper.

Remarks/Arguments begin on page 7 of this paper.

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AMENDMENTS TO THE CLAIMS

1. **(Currently amended)** A method for identifying and/or quantifying an organism or part of an organism in a sample by detecting a nucleotide sequence specific of said organism, wherein said specific nucleotide sequence presents a homology higher than 60% with at least 4 other homologous nucleotide sequences from other organisms comprising:

amplifying ~~or copying~~ said specific nucleotide sequence by PCR into double-stranded target nucleotide sequence using primer pairs which are capable of amplifying at least two of said homologous nucleotide sequences from other organisms;

contacting said target nucleotide sequence with single-stranded capture nucleotide sequences, said single-stranded capture nucleotide sequences being covalently bound in an array to an insoluble solid support via a spacer comprising a nucleotide sequence of at least 40 bases which is at least 6.8 nm in length, wherein said array comprises at least 4 different bound single-stranded capture nucleotide sequences/cm² of solid support surface and wherein said capture nucleotide sequences comprise a nucleotide sequence of about 15 to about 40 bases which is able to specifically bind to said target nucleotide sequence without binding to said at least 4 homologous nucleotide sequences; and

detecting specific hybridization of said target nucleotide sequence to said capture nucleotide sequences.

2. **(Previously presented)** The method according to claim 1, wherein the amplified nucleotide sequence is a DNA nucleotide sequence.

3. **(Canceled)**

4. **(Previously presented)** The method according to claim 1, wherein the amplified nucleotide sequences are mRNA first reverse transcribed into cDNA and then amplified using said primer pair which is capable of amplifying at least two of said homologous mRNA in said sample.

5-7. **(Canceled)**

8. **(Canceled)**

9. **(Previously presented)** The method according to claim 1, wherein the density of the capture nucleotide sequence bound to the surface at a specific location is more than about 10 fmoles per cm² of solid support surface.

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10. **(Previously presented)** The method according to claim 1, wherein the target nucleotide sequence presents a homology with other homologous nucleotide sequences higher than 30%.

11. **(Canceled)**

12. **(Previously presented)** The method according to claim 1, wherein other primers are present-in the amplification step for the amplification of another nucleotide sequence.

13. **(Previously presented)** The method according to claim 1, wherein the insoluble solid support is selected from the group consisting of: glasses, electronic devices, silicon supports, plastic supports, compact discs, filters, filters, gel layers, and metallic supports.

14. **(Previously presented)** The method according to claim 1, wherein the nucleotide sequence to be identified and/or quantified is an RNA sequence submitted to a reverse transcription of its 3' or 5' end by using a consensus primer.

15. **(Previously presented)** The method according to claim 1, wherein the nucleotide sequence to be identified and/or quantified are from the *FemA* gene of Staphylococci species selected from the group consisting of: *S. aureus*, *S. epidermidis*, *S. saprophyticus*, *S. hominis* and *S. haemolyticus*.

16. **(Previously presented)** The method according to claim 1, wherein the solid support also bears capture nucleotide sequences specific of the homologous sequences specific for the binding with the homologous target nucleotide sequence together with a consensus sequence able to bind to said target nucleotide sequence and to said at least 4 homologous nucleotide sequences.

17. **(Original)** The method according to claim 1, wherein the solid support bears capture nucleotide sequences specific for the identification of two or more staphylococcus species together with a consensus sequence for a Staphylococcus genus identification.

18. **(Previously presented)** The method according to claim 1, wherein the sequence to be identified and/or quantified in the sample belongs to the *MAGE* gene family.

19. **(Previously presented)** The method according to claim 1, wherein the sequence to be identified and/or quantified in the sample belongs to the *HLA-A* genes family.

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20. **(Previously presented)** The method according to claim 1, wherein the sequence to be identified and/or quantified in the sample belongs to the dopamine receptors coupled to the protein G genes family.

21. **(Previously presented)** The method according to claim 1, wherein the sequence to be identified and/or quantified in the sample belongs to the choline receptors coupled to the protein G genes family.

22. **(Previously presented)** The method according to claim 1, wherein the sequence to be detected and/or quantified in the sample belongs to the histamine receptors coupled to the protein G genes family.

23. **(Previously presented)** The method according to claim 1, wherein the sequence to be detected and/or quantified in the sample belongs to the cytochrome p450 forms family.

24. **(Withdrawn)** A diagnostic and/or quantification kit which comprises an insoluble solid support upon which single stranded capture nucleotide sequences are bound, said single stranded capture nucleotide sequences containing a sequence of between about 10 and about 60 bases specific for a target nucleotide sequence to be detected and/or quantified and having a total length comprised between about 30 and about 600 bases, said single stranded capture nucleotide sequences being disposed upon the surface of the solid support according to an array with a density of at least 4 single stranded capture nucleotide sequences/cm² of the solid support surface.

25. **(Withdrawn)** The diagnostic kit according to claim 24, wherein the insoluble solid support is selected from the group consisting of glasses, electronic devices, silicon supports, plastic supports, compact discs, gel layers, metallic supports or a mixture thereof.

26. **(Withdrawn)** The diagnostic kit according to claim 24, wherein the capture nucleotide sequences are specific to a target nucleotide sequence to be detected and/or quantified which is specific for a gene selected from the group consisting of *Staphylococcus* species genes, MAGE genes family, HLA-genes family, dopamine, choline or histamine receptors coupled to the protein G genes family, cytochrome P450 forms family or GMO plants family.

27. **(Withdrawn)** The diagnostic kit according to claim 24, comprising biochips, for identification and/or quantification of 5 bacteria species obtained after amplification of one of their DNA sequences with one consensus primer(s) and detection on an array.

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28. **(Withdrawn)** The diagnostic kit according to claim 24, comprising biochips, for identification and/or quantification of bacteria species together with the identification of the bacterial genus obtained after copying and/or amplification of one of their DNA or RNA sequences with one consensus primer(s) and detection on an array.

29. **(Withdrawn)** The diagnostic kit according to claim 24, comprising biochips, for detection and/or quantification of 15 Staphylococcus species obtained after copying and/or amplification of one of their DNA sequences with one consensus primer(s) and detection on an array.

30. **(Withdrawn)** The diagnostic kit according to claim 24, comprising biochips, for detection and/or quantification of 3 or more MAGE genes obtained after copying and/or amplification of one of their DNA or mRNA sequences with one consensus primer(s) and detection on an array.

31. **(Withdrawn)** The diagnostic kit according to claim 24, comprising biochips, for detection and/or quantification of 3 or more HLA-A sequences obtained after copying and/or amplification of one of their mRNA or DNA sequences with one consensus primer(s) and detection on an array.

32. **(Withdrawn)** The diagnostic kit according to claim 24, comprising biochips, for detection and/or quantification of 3 or more gene sequences of receptors coupled to the protein G obtained after copying and/or amplification of one of their mRNA or DNA sequences with one consensus primer(s) and detection on an array.

33. **(Withdrawn)** The diagnostic kit according to claim 32, comprising biochips, for detection and/or quantification of 3 or more gene sequences of dopamine receptors coupled to the protein G obtained after copying and/or amplification of one of their mRNA or DNA sequences with one consensus primer(s) and detection on an array.

34. **(Withdrawn)** The diagnostic kit according to claim 32, comprising biochips, for detection and/or quantification of 3 or more gene sequences of serotonin receptors coupled to the protein G obtained after copying and/or amplification of one of their mRNA or DNA sequences with one consensus primer(s) and detection on an array.

35. **(Withdrawn)** The diagnostic kit according to claim 32, comprising biochips, for detection and/or quantification of 3 or more gene sequences of histamine receptors coupled to the

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protein G obtained after copying and/or amplification of one of their mRNA or DNA sequences with one consensus primer(s) and detection on an array.

36. **(Withdrawn)** The diagnostic kit according to claim 24, comprising biochips, for detection and/or quantification of 3 or more gene sequences of GMO plants obtained after copying and/or amplification of one of their mRNA or DNA sequences with one consensus primer(s) and detection on an array.

37. **(Withdrawn)** The diagnostic kit according to claim 24, comprising biochips, for detection and/or quantification of 3 or more gene sequences the cytochrome P450 forms obtained after copying and/or amplification of one of their mRNA or DNA sequences with one consensus primer(s) and detection on an array.

38. **(Previously presented)** The method of Claim 1, wherein said nucleotide sequence to be identified and/or quantified originates from a microorganism.

39. **(Cancelled)**

40. **(Previously presented)** The method according to claim 1, wherein the density of the capture nucleotide sequence bound to the surface at a specific location is more than about 100 fmoles per cm² of solid support surface.

41. **(Cancelled)**

42. **(Previously presented)** The method according to claim 1, wherein the target nucleotide sequence presents a homology with other homologous nucleotide sequences higher than 80%.

43. **(Cancelled)**

44. **(Previously presented)** The method of Claim 12, wherein said other nucleotide sequence is an antibiotic resistance determining sequence.

45. **(Previously presented)** The method of Claim 1, wherein said organism is identified or quantitated by detecting a single spot signal at one specific location on said insoluble solid support.

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REMARKS

Claim 1 has been amended. Support for the amendment can be found in Examples 3, 4, 7-15 and in paragraph [0044] of the Specification as filed. Therefore, no new matter has been introduced by this amendment. Claim 43 has been canceled without prejudice. The following addresses the substance of the Office Action.

Non-obviousness

The Examiner has rejected Claims 1, 2, 9, 10, 13, 13, 16, 17, 38, 40, and 42-45 under 35 USC §103(a) as being allegedly unpatentable over Anthony et al. (*J. Clin. Microbiol.* 2000 38:781-788) in view of Shchepinov et al. (*Nucleic Acids Res.* 1997 25:1155-1161). More specifically, the Examiner alleges that it would have been obvious to one skilled in the art at the time the invention was made to modify the method of Anthony by using a spacer of at least 6.8 nm in length as taught by Shchepinov et al.

To establish a *prima facie* case of obviousness, the PTO must cite one or more references that provide some suggestion or motivation to modify the references to achieve the claimed invention, provide a reasonable expectation of success to achieve the claimed invention, and finally, the cited art must teach or suggest all the claim limitations. *In re Vaeck*, 947 F.2d 488 (Fed. Cir. 1991). Here, the cited art either taken alone or in combination, fails to provide any of the required factors.

The present invention is related to a method of using arrays comprising covalently bound capture nucleotide sequences wherein these sequences comprise a spacer and a sequence that specifically binds to the target; two categories of capture nucleotides sequences for group and sub-group detection and primer pairs capable of amplifying at least two of 4 homologous sequences. In Claim 1, these single-stranded capture nucleotides sequences are covalently bound to the solid support and include a spacer that places the specific sequence of the capture nucleotide sequence such that it is able to hybridize with the corresponding target nucleotide sequence at a certain distance from the solid support surface (at least 40 bases). The binding between the target nucleotide sequence and its corresponding capture nucleotide sequence forms a signal at the expected location (the location of the specific capture nucleotide sequence), the detection of said signal allowing a discrimination of a target sequence from other homologous sequences obtained from other organisms.

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Furthermore, in contrast to the presently claimed method in which the capture nucleotide sequences are covalently bound to an insoluble support by a spacer, Anthony et al. teaches the use of short capture probes of 20-25 bases which do not include a spacer and are immobilized on nylon membranes. The binding of the capture probes on filter or membrane means that there is no control of which part of the sequence would be available for hybridization (Anthony et al. pg. 783, line 2: "The length of the UV exposure used to link the probe on the nylon was found to have a marked effect on the intensity of the resulting spots"). Therefore, because there is no single point attachment of capture probes on nylon membranes, the addition of a spacer to the capture probes is not compatible with the method of Anthony et al. In contrast to the method of Anthony, the utilization of spacers in the present method ensures that the portions of the capture nucleotide sequences which are complementary to the target sequences are available for hybridization. Therefore, Anthony in fact teaches away from the present invention.

Additionally, in the method of Anthony et al. one target sequence can cross-react with several capture probes (note that some of the filters depicted in Fig. 1 of the Anthony reference contain more than one position of hybridization). In such cases, it is the pattern of several positive spots which allows specific identification of the organism present. This means that the interpretation of the result is not straightforward. Contrary to method which utilize a pattern of spots (i.e. Anthony et al.), in the embodiment of Claim 45 a single "spot signal" directly allows the identification of a specific organism, therefore one capture nucleotide sequence is sufficient for the identification of one target nucleotide sequence thus permitting correlation between intensity of the spot signal and the amount of target nucleotide sequence present.

Combination of references

Shchepinov et al. describes the use of a chemical spacer to reduce steric interference of the support on the hybridization behavior of immobilized oligonucleotides. The support used by Shchepinov et al. is polypropylene which is highly hydrophobic despite amination (see page 1158, left column). If the oligonucleotide is fixed directly onto the surface of a hydrophobic support, there would be strong repulsion during hybridization with a complementary strand. The authors state that negatively charged group in the spacer diminishes the yield of hybridization, and describe the preferred use of a spacer with low negative charge density therefore by stating that high negative charge in the spacer could act to repel the target and hence reduce the rate of

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duplex formation are “teaching away” from the invention in which the spacer is a negatively charged nucleic acid sequence. Furthermore, the authors hybridize with complementary single-stranded short oligonucleotides (12 nucleotides) or with tRNA which is also single-stranded. Additionally, the optimal chemical spacer is 30-60 atoms in length which would be equivalent to 15-30 nucleotides in length (see figure 2). Duplex yield declines with further increase in the length of the spacer. At 30 units (180 atoms or 90 nucleotides), the yield of hybridization is the same as that with no spacer at all (figure 3b).

In the present invention, the inventors have used a spacer made of nucleotides sequences of at least 40 nucleotides which are negatively charged (at least 40 negative charges). The yield of hybridization is still increasing with a spacer of 60 nucleotides (see Figure 3 as filed). The unexpected advantage of the method of the present invention is that a very good yield of hybridization of amplified long target DNA was obtained even in the presence of its complementary nucleotide sequence present in solution. The hybridization is performed with double-stranded amplified by PCR and present in solution target DNA. The dsDNA reassociate much faster in solution than to hybridize on small sequences fixed on a solid support where diffusion is low, thus reduce even more the rate of the reaction. The difference of the claimed method is the use of small specific sequences on capture probes in order to be able to differentiate between homologous sequences while having a high yield of hybridization on the immobilized probe (see paragraph [0044] of the Specification as filed).

Shchepinov et al. neither mention nor suggest the use of spacer of at least 40 nucleotides. In fact, the optimal spacer length of Shchepinov et al. (equivalent to 15-30 nucleotides) is not within the range of the present invention (above 40 nucleotides). As shown in the figure 3 of the Specification as filed, the rate of hybridization is increased by a factor of ~2 for a spacer of 60 nucleotides as compared to a spacer of 20 nucleotides while there is no increase at this size in the paper of Shchepinov et al.

Additionally, Shchepinov et al. method is not related to a detection of homologous sequences contrary to the present invention. The solution proposed by the present invention is to obtain both a specific and sensitive detection of multiple homologous sequences which are present in solution as amplified double-stranded nucleotide sequences. Hybridization conditions are much more stringent in case of homologous sequences in order to differentiate small

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differences in their sequences, which still reduce the hybridization rate. Results are therefore hardly comparable.

The high yield of hybridization is an important aspect of the present invention since it allows the surface bound capture probes to efficiently compete with the reassociation of the target with its complementary sequence present in the same solution. The inventors found that a spacer of at least 40 bases increases the specificity of hybridization of the capture molecule with the target molecule. The spacers of Shchepinov et al. are built up from monomeric units, using phosphoramidite chemistry by condensation onto an amine functionalized polypropylene support. This means that they are synthesized directly on the support according to Southern's methods. Long polynucleotides including a spacer of more than 40 nucleotides and a specific sequence of 15-40 nucleotides cannot be directly synthesized on the support due to limits of the *in situ* synthesis. Indeed, current coupling efficiencies impose a limit of \pm 25 bases to these chips. Beyond this limit, the accumulation of incomplete synthesis products becomes a problem. The spacer of the present invention is not synthesized directly on the support but by conventional chemical synthesis. The specific part of the capture probe may be synthesized consecutively to the spacer nucleotide bases resulting in a single molecule. The synthesized capture nucleotide sequences are then covalently fixed on the support according to an array by physical deposition. The single molecules thus obtained highly simplify and reduce the cost of production of the capture nucleotide sequences.

For these reasons, it would not have been obvious to one of ordinary skill in the art at the time the invention was made to have modified the method of Anthony et al. so as to have used a chemical spacer of Shchepinov et al. These references both fail because neither provides the requisite motivation to combine, the reasonable expectation of success, nor teach all the limitations of the claimed invention.

Furthermore, the inventors completed the invention of this application prior to February 2000, the date that appears on Anthony et al. publication, and therefore the cited reference does not constitute prior art. The Declaration under 37 CFR §1.131 supporting this assertion will be submitted shortly.

Therefore, Claims 1, 2, 9, 10, 13, 13, 16, 17, 38, 40, and 42-45 are not obvious over the cited references and their rejection under 35 USC §103(a) should be withdrawn.

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The Examiner has rejected Claim 15 under 35 USC §103(a) as being allegedly unpatentable over Anthony et al. (*J. Clin. Microbiol.* 2000 **38**:781-788) in view of Shchepinov et al. (*Nucleic Acids Res.* 1997 **25**:1155-1161) as applied to Claims 1, 2, 9, 10, 13, 13, 16, 17, 38, 40, and 42-45 above and further in view of Vannuffel et al. (WO 99/16780). More specifically, the Examiner alleges that because Vannuffel et al. teaches the detection of the *FemA* gene of *Staphylococci* species, it would have been obvious to combine this teaching with the method of Anthony et al. as modified by using a spacer of at least 6.8 nm in length as taught by Shchepinov et al.

Vannuffel's disclosure of the use of the *FemA* gene of *Staphylococci* species to detect bacteria present in a sample does not provide motivation to combine arrays comprising covalently bound probes of the lengths recited in the claims which comprise a spacer of at least 40 nucleotides in length, and primer pairs capable of amplifying at least two of 4 homologous sequences in methods for detecting specific nucleotide sequences having a homology level higher than 60% with sequences from other organisms. Furthermore, Vannuffel fails to cure the deficiencies of Anthony et al. combined with Shchepinov et al. as discussed with regards to Claims 1, 2, 9, 10, 13, 13, 16, 17, 38, 40, and 42-45 above. Thus, Vannuffel et al. fails to correct the failure of Anthony et al., and Shchepinov et al. to render the claimed invention obvious for the reasons addressed above. Therefore, dependent Claim 15 is also non-obvious.

The Examiner has rejected Claim 18 under 35 USC §103(a) as being allegedly unpatentable over Anthony et al. (*J. Clin. Microbiol.* 2000 **38**:781-788) in view of Shchepinov et al. (*Nucleic Acids Res.* 1997 **25**:1155-1161) as applied to Claims 1, 2, 9, 10, 13, 13, 16, 17, 38, 40, and 42-45 above and further in view of Boon et al. (USP 6,488,932). More specifically, the Examiner alleges that because Boon et al. teach that it is advantageous to detect individual sequences that belong to MAGE family for the diagnosis of tumors, it would have been obvious to combine this teaching with the method of Anthony et al. as modified by using a spacer of at least 6.8 nm in length as taught by Shchepinov et al.

Boon's disclosure of the use of the MAGE family to diagnose tumors does not provide motivation to combine arrays comprising covalently bound probes of the lengths recited in the claims which comprise a spacer of at least 40 nucleotides in length, and primer pairs capable of amplifying at least two of 4 homologous sequences in methods for detecting specific nucleotide

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sequences having a homology level higher than 60% with sequences from other organisms. As discussed above, Anthony et al. and Shchepinov et al. do not disclose all the limitations of the claimed invention, and Boon *et al.* fails to correct the failure of Anthony *et al.*, and Shchepinov *et al.* to render the claimed invention obvious for the reasons addressed above. Therefore, dependent Claim 18 is also non-obvious.

The Examiner has rejected Claims 4, 14 and 19 under 35 USC §103(a) as being allegedly unpatentable over Anthony et al. (*J. Clin. Microbiol.* 2000 38:781-788) in view of Shchepinov et al. (*Nucleic Acids Res.* 1997 25:1155-1161) as applied to Claims 1, 2, 9, 10, 13, 13, 16, 17, 38, 40, and 42-45 above and further in view of Apple et al. (USP 5,451,512). More specifically, the Examiner alleges that because Apple et al. teach that it is advantageous to detect individual sequences that belong to HLA-A family to help determine potential transplantation donors and teaches the amplified nucleotide sequences for HLA-A detection, it would have been obvious to combine this teaching with the method of Anthony et al. as modified by using a spacer of at least 6.8 nm in length as taught by Shchepinov et al.

Apple's disclosure of the use of HLA-A family to help determine potential transplantation donors does not provide motivation to combine arrays comprising covalently bound probes of the lengths recited in the claims which comprise a spacer of at least 40 nucleotides in length, and primer pairs capable of amplifying at least two of 4 homologous sequences in methods for detecting specific nucleotide sequences having a homology level higher than 60% with sequences from other organisms. As discussed above, Anthony et al. and Shchepinov et al. do not disclose all the limitations of the claimed invention, and Apple *et al.* fails to correct the failure of Anthony *et al.*, and Shchepinov *et al.* to render the claimed invention obvious for the reasons addressed above. Therefore, dependent Claims 4, 14 and 19 are also non-obvious.

The Examiner has rejected Claims 20 and 22 under 35 USC §103(a) as being allegedly unpatentable over Anthony et al. (*J. Clin. Microbiol.* 2000 38:781-788) in view of Shchepinov et al. (*Nucleic Acids Res.* 1997 25:1155-1161) as applied to Claims 1, 2, 9, 10, 13, 13, 16, 17, 38, 40, and 42-45 above and further in view of Klein et al. (USP 6,255,059). More specifically, the Examiner alleges that because Klein teaches that it is advantageous to detect sequences that belong to the dopamine or histamine receptors coupled to the G genes family to mediate

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transmembrane signaling by external stimuli, endocrine function, carbohydrate metabolism, etc., it would have been obvious to combine this teaching with the method of Anthony et al. as modified by using a spacer of at least 6.8 nm in length as taught by Shchepinov et al.

Klein's disclosure of the use of sequences that belong to the dopamine or histamine receptors coupled to the G genes family to mediate transmembrane signaling by external stimuli, endocrine function, carbohydrate metabolism, etc. does not provide motivation to combine arrays comprising covalently bound probes of the lengths recited in the claims which comprise a spacer of at least 40 nucleotides in length, and primer pairs capable of amplifying at least two of 4 homologous sequences in methods for detecting specific nucleotide sequences having a homology level higher than 60% with sequences from other organisms. As discussed above, Anthony et al. and Shchepinov et al. do not disclose all the limitations of the claimed invention, and Klein et al. fails to correct the failure of Anthony et al., and Shchepinov et al. to render the claimed invention obvious for the reasons addressed above. Therefore, dependent Claims 20 and 22 are also non-obvious.

The Examiner has rejected Claim 21 under 35 USC §103(a) as being allegedly unpatentable over Anthony et al. (*J. Clin. Microbiol.* 2000 38:781-788) in view of Shchepinov et al. (*Nucleic Acids Res.* 1997 25:1155-1161) as applied to Claims 1, 2, 9, 10, 13, 13, 16, 17, 38, 40, and 42-45 above and further in view of Murphy et al. (WO 94/05695). More specifically, the Examiner alleges that because Murphy teaches that it is advantageous to detect sequences that belong to the choline receptors coupled to the G genes family for use in diagnosis of neurological, viral or endocrine pathologies, it would have been obvious to combine this teaching with the method of Anthony et al. as modified by using a spacer of at least 6.8 nm in length as taught by Shchepinov et al.

Murphy's disclosure of the use of sequences that belong to the choline receptors coupled to the G genes family for use in diagnosis of neurological, viral or endocrine pathologies does not provide motivation to combine arrays comprising covalently bound probes of the lengths recited in the claims which comprise a spacer of at least 40 nucleotides in length, and primer pairs capable of amplifying at least two of 4 homologous sequences in methods for detecting specific nucleotide sequences having a homology level higher than 60% with sequences from other organisms. As discussed above, Anthony et al. and Shchepinov et al. do not disclose all the

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limitations of the claimed invention, and Murphy *et al.* fails to correct the failure of Anthony *et al.*, and Shchepinov *et al.* to render the claimed invention obvious for the reasons addressed above. Therefore, dependent Claim 21 is also non-obvious.

The Examiner has rejected Claim 23 under 35 USC §103(a) as being allegedly unpatentable over Anthony et al. (*J. Clin. Microbiol.* 2000 38:781-788) in view of Shchepinov et al. (*Nucleic Acids Res.* 1997 25:1155-1161) as applied to Claims 1, 2, 9, 10, 13, 13, 16, 17, 38, 40, and 42-45 above and further in view of Waxman et al. (USP 6,207,648). More specifically, the Examiner alleges that because Waxman teaches that it is advantageous to detect sequences that belong to the cytochrome P450 isoforms family for use in treatment of cancer, it would have been obvious to combine this teaching with the method of Anthony et al. as modified by using a spacer of at least 6.8 nm in length as taught by Shchepinov et al.

Waxman's disclosure of the use of sequences that belong to the cytochrome P450 isoforms family for use in treatment of cancer does not provide motivation to combine arrays comprising covalently bound probes of the lengths recited in the claims which comprise a spacer of at least 40 nucleotides in length, and primer pairs capable of amplifying at least two of 4 homologous sequences in methods for detecting specific nucleotide sequences having a homology level higher than 60% with sequences from other organisms. As discussed above, Anthony et al. and Shchepinov et al. do not disclose all the limitations of the claimed invention, and Waxman *et al.* fails to correct the failure of Anthony *et al.*, and Shchepinov *et al.* to render the claimed invention obvious for the reasons addressed above. Therefore, dependent Claim 23 is also non-obvious.

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CONCLUSION

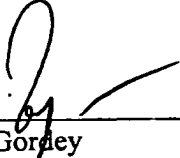
In view of the foregoing, Applicants respectfully submit the present application is fully in condition for allowance. If any issues remain that may be addressed by a phone conversation, the Examiner is invited to contact the undersigned at the phone number listed below.

Please charge any additional fees, including any fees for additional extension of time, or credit overpayment to Deposit Account No. 11-1410.

Respectfully submitted,

KNOBBE, MARTENS, OLSON & BEAR, LLP

Dated: September 19, 2005

By: 

Marina L. Gordey
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INFORMATION DISCLOSURE STATEMENT

Applicant : Remacle et al.
App. No : 09/817,014
Filed : March 23, 2001
For : IDENTIFICATION OF BIOLOGICAL
(MICRO)ORGANISMS BY
DETECTION OF THEIR
HOMOLOGOUS NUCLEOTIDE
SEQUENCES ON ARRAYS
Examiner : Calamita, Heather
Art Unit : 1637

Mail Stop Amendment
Commissioner for Patents
P.O. Box 1450
Alexandria, VA 22313-1450

Dear Sir:

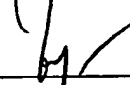
Enclosed for filing in the above-identified application is a PTO/SB/08 Equivalent listing 3 references to be considered by the Examiner. Also enclosed are 3 foreign patent references and/or non-patent literature as listed on the Information Disclosure Statement.

This Information Disclosure Statement is being filed before the mailing date of a final action and before the mailing of a Notice of Allowance. This Statement is accompanied by the fees set forth in 37 C.F.R. § 1.17(p). The Commissioner is hereby authorized to charge any additional fees which may be required or to credit any overpayment to Account No. 11-1410.

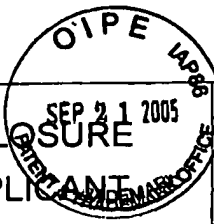
Respectfully submitted,

KNOBBE, MARTENS, OLSON & BEAR, LLP

Dated: September 19, 2005

By: _____ 

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PTO/SB/08 Equivalent

INFORMATION DISCLOSURE STATEMENT BY APPLICANT		Application No.	09/817,014
		Filing Date	March 23, 2001
		First Named Inventor	Remacle et al.
		Art Unit	1637
(Multiple sheets used when necessary)		Examiner	Calamita, Heather
SHEET 1 OF 1		Attorney Docket No.	VANM213.001AUS

U.S. PATENT DOCUMENTS

FOREIGN PATENT DOCUMENTS

FOREIGN PATENT DOCUMENTS						
Examiner Initials	Cite No.	Foreign Patent Document Country Code-Number-Kind Code Example: JP 1234567 A1	Publication Date MM-DD-YYYY	Name of Patentee or Applicant	Pages, Columns, Lines Where Relevant Passages or Relevant Figures Appear	T ¹

NON PATENT LITERATURE DOCUMENTS

Examiner Initials	Cite No.	Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published.	T ¹
	1	OFFICE ACTION from co-pending U.S. Patent Application No. 10/056,229, dated February 25, 2003 (VANM213.001CP1).	
	2	OFFICE ACTION from co-pending U.S. Patent Application No. 10/056,229, dated January 2, 2004 (VANM213.001CP1).	
	3	Letter from Jose Remacle to Eric Van Maideren, dated February 24, 2000.	

1924478:vr091205

***Examiner:** Initial if reference considered, whether or not citation is in conformance with MPEP 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to applicant.

T¹ - Place a check mark in this area when an English language Translation is attached.